

The Microbial Detection Array Combined with Random Phi29-Amplification used as a Diagnostic Tool for Unbiased Virus Detection in Clinical Samples

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ABSTRACT

A common technique used for sensitive and specific diagnostic virus detection in clinical samples is PCR. However, an unbiased diagnostic microarray containing probes for all human pathogens could replace hundreds of individual PCR-reactions and thereby remove the need for a clear clinical hypothesis regarding the suspected pathogen. We have established such a diagnostic platform for unbiased random amplification and microarray identification of viral pathogens in clinical samples. We show that Phi29 polymerase-amplification of a diverse set of clinical samples generates enough viral material for successful identification by the Microbial Detection Array developed at the Lawrence Livermore National Laboratory, California, USA, demonstrating the potential of the microarray technique for broad-spectrum pathogen detection. We conclude that this method detects both DNA and RNA virus, present in the same sample, as well as differentiates between different subtypes. We propose this assay for unbiased diagnostic analysis of all viruses in clinical samples.

INTRODUCTION

A common method of choice for clinical pathogen identification is polymerase chain reaction (PCR) [1] which is a sensitive and specific method. However, each PCR reaction only provides identification of one specific virus, or a group of related viruses. Thus, a clinical hypothesis regarding the suspected pathogen needs to guide the decision as to which PCR analyses to perform, resulting in a large number of PCR protocols needed to facilitate the identification of all human pathogens. Moreover, with some clinical syndromes or symptoms, it might be difficult to determine which viruses to test for in order to establish the correct diagnosis.

DNA microarray testing has emerged as a promising new technology for broad-spectrum virus detection, making it possible to test for the presence of thousands of viruses simultaneously. Several microarray platforms have been developed for detection of all known viruses, as well as novel viruses related to known virus families [2,3,4]. Some of them have been tested on clinical samples, mostly respiratory samples [2,3,4,5]. However, without a pre-amplification of the clinical sample, such as PCR or virus culture, most microarrays are not sensitive enough for pathogen identification since virus titres normally are below their detection limit. For example, a diagnostic human Papilloma virus (HPV)-specific microarray utilizes an HPV-specific PCR-amplification prior to microarray hybridization [6]. Different PCR-based random amplifications have been developed to achieve enough material for a successful identification [3,4,7]. However, studies have shown that PCR-based whole genome amplification (WGA) may generate incomplete coverage of genomes [8]. Instead we used multiple displacement amplification (MDA)-based WGA through isothermal amplification by the Phi29 polymerase [8]. The Phi29 polymerase has the highest processivity rate reported, ~70 000 bases every time it binds [9] and an error rate of only 1 in 10^6 - 10^7 bases [10],

allowing for good coverage across genomes [8,11,12]. The average length produced is >10 kb compared to ≤ 3 kb by the Taq polymerase [8]. Phi29-amplification prior to microarray analysis has been used for human genomic DNA analysis [13,14,15] and for bacterial DNA analysis [16,17]. However, the Phi29 polymerase cannot amplify RNA or short DNA fragments (<2000 bp), such as cDNA generated with random hexamers. To overcome this, the method Whole Transcriptome Amplification (WTA) has included a ligation step prior to amplification [2] resulting in cDNA fragments that are ligated into larger transcriptomes that can be efficiently amplified by the Phi29 polymerase.

Here we investigate real life utility and sensitivity of the microarray technique in a diagnostic set-up. We have established a protocol including pre-treatment, nucleic acid (NA) purification and Phi29-amplification followed by microarray analysis of DNA and RNA viruses from a diverse set of clinical samples. We show that Phi29-amplification generates enough material for successful identification by the Microbial Detection Array (MDA) [18,19], demonstrating the potential of the microarray technique for broad-spectrum pathogen detection in clinical samples. Furthermore, we can detect both DNA and RNA viruses simultaneously present in a sample, as well as differentiate between different subtypes. We propose this assay for unbiased diagnostic analysis of all viruses in clinical samples, helping clinicians in determining the correct diagnosis.

MATERIALS AND METHODS

Virus and clinical samples

This study used clinical samples received for routine diagnostic analysis at the Department of Virology, Statens Serum Institut (SSI), Copenhagen, Denmark (Danish National reference laboratory (ISO 17025); www.ssi.dk), containing DNA or RNA viruses. DNA viral genomes were linear or circular double-stranded: Herpes Simplex virus (HSV) 1 and HSV2, BK Polyomavirus (BKV), JC Polyomavirus (JCV) and HPV type 6, 16, 53, 61. RNA viral genomes were (i) segmented double-stranded: Rotavirus A (Rota A), (ii) (+)single-stranded: Astrovirus, Sapovirus, Dengue 1, Hepatitis C virus (HCV) and Enterovirus A (HEV-A), and (iii) (-)single-stranded: human Respiratory Syncytial virus (RSV). Clinical samples were cerebrospinal fluid (CSF), urine, serum, cervical smear, faeces, skin lesion and tracheal aspirate (TA). Techniques used for routine identification of viruses were in-house real-time PCR assays (for HSV1, HSV2, BKV, JCV, Rota A, Astrovirus, Sapovirus, Dengue, HCV, HEV-A and RSV) or Clart® HPV2 microarray (Genomica) (for HPV). PCR's were performed on Mx3005P thermal cyclers (Stratagene).

Ethics statement

Exemption for review by the ethical committee system and for obtaining informed consent was obtained from the Committee on Biomedical Research Ethics for the Capital region in accordance with Danish law on quality control and assay development projects.

Pre-treatment and extraction of clinical samples

Five- to two-hundred microliters of clinical sample was centrifuged at 17 000 g for 10 minutes. The supernatant was filtered through a 0.22 µm Ultrafree MC spin filter (Millipore) at 2000 g for 2

minutes and DNase treated (6 U DNaseI Amplification grade, Invitrogen) for 1½ hours at room temperature while shaken in a Thermomixer (AH Diagnostics). The viral NA was extracted using the PureLink Viral RNA/DNA kit (Invitrogen), without the addition of carrier RNA. The extracted viral NA was eluted with 20-30 µl DNase/RNase-free sterile water, and stored at -20 °C or immediately used.

Reverse transcription

Reverse transcription on purified viral RNA was performed using the Superscript VILO cDNA synthesis kit (Invitrogen) according to protocol. Briefly, 14 µl of extracted viral RNA was put into a 20 µl-reaction containing Superscript enzyme and random primers, and incubated at 42 °C for 1 hour. The enzyme was inactivated at 95 °C for 5 minutes and the sample stored at -20 °C or immediately used.

Phi29 polymerase amplification

For data presented in Figure 1, viral DNA was amplified by Phi29 polymerase amplification using GenomiPhi V2 Amplification kit (GE Healthcare). Briefly, 1 µl of purified DNA is added to 9 µl of sample buffer, incubated at 95 °C for 3 minutes and cooled on ice. Ten microliters of GenomiPhi reaction is added, containing Phi29 polymerase and random primers, and the reaction incubated at 30 °C for 1 ½ hours. The reaction was terminated by incubation at 65 °C for 10 minutes. The amplified viral DNA was stored at -80 °C or immediately used. For data presented in Figure 4, viral DNA was amplified by Phi29 polymerase amplification using Repli-g Midi kit (typical yield ~40 µg from a 50 µl reaction, Qiagen), following manufacturer's protocols. Briefly, 5 µl of purified DNA is mixed with 5 µl denaturation buffer, incubated at room temperature for 3 minutes and the reaction terminated by the addition of 10 µl stop solution. Thirty microliters of Repli-g reaction

mix, containing Phi29 and random primers, was added and the reaction incubated at 30 °C for 16 hours. The reaction was terminated by incubation at 65 °C for 3 minutes and the amplified viral DNA stored at -80 °C or immediately used.

Whole transcriptome amplification (WTA)

For viral NA amplification (both DNA and RNA virus) we used the described WTA method [2] by using the QuantiTect Whole Transcriptome Amplification kit (typical yield ~40 µg from a 50 µl reaction, Qiagen) according to manufacturer's protocol, except for the reverse transcription step that was replaced by the Superscript VILO kit (Invitrogen) described above. The WTA protocol includes 3 sequential reactions: firstly a reverse transcription reaction to generate cDNA, secondly a ligation of generated cDNA into larger transcriptomes and thirdly a Repli-g amplification of the generated transcriptomes. Briefly, as described above, 14 µl of extracted viral NA was put into a 20 µl-VILO reaction and incubated at 42 °C for 1 hour. The Superscript enzyme was inactivated at 95 °C for 5 minutes, 10 µl of the VILO cDNA reaction was added to 10 µl of ligation mix and incubated at 22 °C for 2 hours. Thirty microliters of Repli-g amplification mix containing Phi29 polymerase and random primers was then added and amplification performed at 30 °C for 8 hours. The reaction was terminated by incubation at 95 °C for 5 minutes. Amplified material was purified using QIAamp DNA mini kit (Qiagen) using a modified protocol (www.qiagen.com), and checked for purity and concentration using a NanoDrop spectrophotometer (Thermo Scientific) and agarose gel electrophoresis. The DNA was then stored at -80 °C or immediately used for microarray analysis.

Quantification and conformation by real-time PCR

For the quantification of virus before and after amplification, virus-specific real-time PCR's were performed. These were in-house assays for HSV1, HSV2, HPV6, HPV16, BKV, JCV, Rota A, Astrovirus, Sapovirus, HCV, HEV-A, RSV and β -actin (detecting human gDNA). In addition, a previously published assay was used for JCV [20]. For the multi-HPV (6, 16, 53, 61) positive sample (determined in the routine by Clart® HPV2 microarray, Genomica), only HPV 6- and 16-specific PCR's were performed before and after Phi29-amplification. For the quantitative analysis of BKV, we used an in-house standard containing 20 000 – 2×10^7 viral copies/ml. For the confirmation of additional viruses found by microarray analysis, in-house real-time PCR's were used for HPV103, Picobirnavirus, Rota C and Dengue subtypes (1, 2, 3 and 4), and previously published assays for JCV [20], GB virus type C (GBV-C) [21] and Human Adenovirus C (HAdV-C= [22]. Thermal cycling was performed in a thermal cycler (Mx3005P (Stratagene) or ABI7900 (Applied Biosystems)). Fold difference was calculated from ΔC_t obtained from real-time PCR results before and after Phi29-amplification, combined with dilution factors for each sample. In doing so, we also made the assumption that a change in C_t -value was equivalent to a doubling of target DNA.

Microarray analysis

The microarray used was the MDA version 2, developed at Lawrence Livermore National Laboratory (LLNL), USA and already described elsewhere [18,19]. Briefly, the MDA contains 388,000 probes designed from all sequenced virus and bacteria, resulting in a large number of probes covering each genome. The microarray was designed for both detection and discovery, to both identify known organisms and to detect un-sequenced or emerging organisms, while strain or subtype identification was not a goal. The array have both conserved and unique probes within each

family, where oligos that were conserved, to the extent possible, within a family, and unique relative to other families and kingdoms, were prioritized [18]. Labelling and microarray hybridization was performed at NimbleGen Systems-Iceland, Reykjavik, Iceland, or in-house at SSI, Copenhagen, Denmark according to manufacturer's protocol (Gene expression analysis, Roche NimbleGen). Briefly, 1 µg of heat-denatured WTA-amplified sample was labelled using nick translation with Cy3-labeled random nonamer primers (TriLink Biotechnologies) and Klenow fragment (exo-) (NEB) at 37 °C for 2 hours. Labelled DNA was iso-propanol precipitated and the pellet washed, dried, reconstituted and quantitated. For hybridization, the in-house assay used 12 µg of labelled DNA whereas NimbleGen Systems-Iceland used 6 µg. Hybridization was performed in a NimbleGen Hybridization system using the NimbleGen Hybridization kit (Roche NimbleGen).

Microarrays were scanned using a GenePix 4000B Scanner (Molecular Devices) and data analyzed using the maximum likelihood method developed at LLNL, extensively described elsewhere [18]. Additional stringency criteria were applied to exclude bacterial sequences and viruses having fewer than 20% of probes detected out of those expected. Briefly, probes were classified as detected if the intensity exceeded a threshold equal to the 99th percentile of intensities for negative control probes, except that one sample was analyzed using the 95th percentile. Targets in an internal database of 43 705 viral sequences, developed at LLNL [18], were screened against the stringency criteria; an unconditional log-odds score for presence of each remaining target was computed, and targets having log-odds scores less than 5 were excluded. A greedy forward selection algorithm was applied to find the collection of targets most likely to be present in the sample. At every forward selection step, a conditional log-odds score was computed for each remaining target, representing its potential contribution to the log-likelihood conditioned on the presence of the previously selected targets; the target having the largest conditional log-odds was selected and added to the

collection. The resulting conditional and unconditional log-odds scores were plotted in a bar graph format [18].

As a complement, another mode of data analysis was developed at SSI, where data was processed using the NimbleScan software program (NimbleGen, Roche). A mean of intensities for each probe-set was calculated. Outlier intensities within each probe-set were filtered out when signals were more than 3 standard deviations from the mean of the probe-set. A signal threshold was defined as the 99th percentile of the random control intensities, and every probe-set mean containing more than 10 signals above this threshold was considered as a positive signal.

RESULTS

Sample pre-treatment and Phi29-amplification

To remove contaminating human gDNA present in clinical samples, we designed a pre-treatment protocol including centrifugation, filtration and DNase treatment. It has been demonstrated that filtration, DNase and RNase treatment can increase the efficiency of downstream amplifications [23,24]. During DNase treatment the viral NA should be protected inside the viral particle [25]. After pre-treatment, the viral particles were lysed, viral NA purified and later Phi29-amplified by using GenomiPhi (GE Healthcare). During purification, the addition of carrier RNA did not improve the yield and interfered with the downstream Phi29-amplification, and was left out (data not shown). Quantitative analysis of viral DNA or human gDNA before and after pre-treatment or Phi29-amplification was done by real-time PCR. Only centrifugation and filtration left large amounts of human gDNA still present in a HSV1⁺ sample (Ct = 22), while DNase treatment reduced the β -actin signal 1000-fold (Ct = 32) (Figure 1A). To investigate to what extent Phi29-amplification of remaining human gDNA would occur we performed a time-course experiment with the amplification-time extended up to 16 hours (Figure 1B). The β -actin signal was still at acceptable background level (3000-fold increase in signal), while the HSV1 signal showed an 80 000-100 000-fold increase in signal. In conclusion, our pre-treatment removed a significant amount of human gDNA without losing the virus-specific signal, allowing for good virus amplification.

Whole transcriptome amplification

To establish an amplification protocol suitable for both DNA and RNA viruses, we used the described Phi29-amplification method WTA [2] that has included a ligation step prior to the amplification step, thereby generating larger templates for Phi29. The WTA performed on a HCV⁺

serum sample (RNA virus) resulted in a ~200 000-fold increase in signal (Figure 2A and Table 1), as compared to only 3-fold without ligation (data not shown). The WTA RT-reaction and ligation step did not have any adverse effects when performed on a DNA-virus such as JCV from a CSF sample (Figure 2B and Table 1), enabling us to run both viral DNA and RNA through the same amplification protocol prior to microarray hybridization. The WTA was used on 12 different DNA or RNA viruses found in 7 different types of clinical samples (Table 1 and Figure 2). The fold increase in PCR signal comparing before and after WTA, ranged from 330- to 564×10^6 -fold. The highest fold increase was detected for the circular viruses JCV, BKV and HPV (Table 1 and Figure 2B). Such circular genomes have previously been shown to be efficiently amplified by Phi29 polymerase, through rolling circle amplification (RCA) [26,27]. The lowest increase in signal was seen for Rota A found in faeces (Table 1) due to large quantities already present before amplification ($C_t=12$), which resulted in a C_t -value of 7 after WTA amplification (Table 1). WTA-amplified samples were purified and typical yield after 8 hours of WTA was ~25 µg (data not shown) showing good size distribution (Figure 2C).

Microarray analysis of clinical samples

Samples amplified by WTA representing 7 different types of clinical samples were labelled and hybridized to the MDA (Figure 3 and Table 2). Microarray data was analysed using the maximum likelihood method developed at LLNL [18,19], with additional stringencies applied. In 13 out of 13 samples, the expected virus was detected: HSV1, HSV2, HPV16, HPV6/16/53/61, BKV, JCV, Rota A, Astrovirus, Sapovirus, Dengue 1, HCV in duplicate, HEV-A and RSV (Table 2). For the RSV⁺ sample, a detection threshold equal to the 95th percentile of control probes had to be used for detection. For this sample we also used the data analysis developed at SSI, where RSV was detected using a detection threshold equal to the 99th percentile of random control probes (data not shown).

Moreover, using this approach on all samples, we generated the same result as seen with the maximum likelihood method (data not shown). For the double-positive faeces sample containing both Astro- and Sapovirus, only Astrovirus was detected by the microarray analysis. In the negative control no viruses were detected.

In 10 out of 13 clinical samples (77 %) representing all 7 types of clinical samples used in the study (skin lesion, urine, CSF, cervical smear, serum, faeces and TA), viruses from the Anelloviridae family (circular ssDNA) were found (Torque Teno virus (TTV), Torque Teno Midi virus (TTMV) and TTV-like mini virus) (Table 2). They were detected in samples containing DNA or RNA viruses, demonstrating that our pre-treatment and amplification protocol followed by microarray analysis can easily detect both types in one sample. Additional viruses were detected in 9 samples (Table 2), with log-odds scores ranging from 91.0 to 677.5. Some were clinically irrelevant since they were phages (Propionibacterium phage, Phage Gifsy-2, Lactococcus phage, Enterobacteria phage ST104, Mycobacteriophage and Pseudomonas phage) or plant viruses (Pepino mosaic virus), probably from digested or environmental sources. Human endogenous retrovirus (HERV) was detected in the Dengue 1 and the RSV sample (Table 2). The rest of the additional viruses found (HPV103, Picobirnavirus, HAdV-C, GBV-C, JCV, Rota C and Dengue 2) were tested with virus-specific PCR for confirmation. The HPV103 (cir dsDNA) detected in the HPV16⁺ sample is not present on the microarray used for routine HPV diagnosis (Genomica) instead its presence was confirmed using a HPV103-specific PCR (data not shown). The Rota A⁺ faeces sample was found to be positive for Picobirnavirus (dsRNA) and negative for Rota C (dsRNA) by PCR (data not shown). The Sapovirus⁺ faeces sample was found to be positive for HAdV-C (linear dsDNA) in the microarray analysis, which was confirmed by PCR. Two separate microarray experiments detected GBV-C ((+)ssRNA) in the HCV⁺ sample, which was confirmed by a GBV-C-specific PCR (data

not shown). The BKV⁺ sample was detected as positive for JCV by microarray but tested negative by PCR (data not shown). The Dengue 1⁺ sample was found to be positive for both Dengue-1 and Dengue 2 in the microarray analysis, but tested negative for types 2, 3 and 4 using Dengue subtype-specific PCR's (data not shown).

Microarray detection range for BKV⁺ urine samples

To investigate the detection range of the microarray on clinical samples, we performed analysis on urine samples positive for the circular virus BKV (Table 3 and Figure 4). Samples ranging from 15 810 to 4.0×10^8 copies/ml of BKV were amplified by Phi29 polymerase for 16 hours. Also included were three dilutions of a BKV urine sample, containing 892, 119 or 73 copies/ml, respectively. Samples containing ≥ 1000 copies/ml of BKV (BKV1-BKV5) were efficiently amplified with yields from 1.0×10^{11} to 1.9×10^8 copies/ml of BKV (Table 3 and Figure 4A), and with ~ 25 μ g of DNA generated after amplification (data not shown). The total input of viral copies per 50 μ l Phi29-reaction for BKV1-BKV5, were 2×10^6 , 4100, 2100, 80 and 5 genomic copies, respectively (Table 3). Amplification of samples containing ~ 100 copies/ml (BKV6 and BKV7) resulted in loss of reproducibility and low yields (Table 3 and Figure 4A). The total input of viral copies per Phi29-reaction for these two samples were 0.6 and 0.4, respectively (Table 3). Theoretically, amplification of samples containing 200 copies/ml means that ~ 1 viral copy is added to the Phi29-amplification reaction, resulting in stochastic problems due to unequal distribution in highly diluted solutions. Four amplifications were selected for microarray analysis (BKV3, BKV4, BKV5 and BKV6) and hybridized to the MDA (Table 3 and Figure 4B). BKV⁺ urine samples containing ≥ 1000 copies/ml were clearly detected by the microarray analysis after Phi29-amplification (Figure 4B). The human genomic sequence SSX3 (synovial sarcoma, X breakpoint 3) was used as a hybridization control and had the same signal intensity in all 4 microarrays. Some cross-reactivity with JCV was seen in

the 3 samples where BKV was detected, even if they all were negative in a JCV-specific PCR (data not shown). This is similar to the BKV sample presented in table 2. From this we conclude that, for a circular genome and from as little as 5 copies input (≥ 1000 copies/ml) the Phi29-amplification was able to generate enough material for a clear detection by microarray analysis.

DISCUSSION

By using WTA for viral amplification, both DNA and RNA viruses could be run through the same protocol. The correct DNA or RNA virus was detected in 13 out of 13 samples, representing a diverse set of clinical materials. Moreover, we can correctly detect multiple subtypes of a virus present in a sample, as seen for two clinical samples positive for multiple HPV types (types 6, 16, 53 and 61, and types 16 and 103, respectively), with no cross-reactivity towards other subtypes. It should be noted that sub-type identification was not a goal in the probe design when developing MDA, nevertheless, the ability to combine the signals from multiple probes in the analysis made it possible to discriminate between different HPV subtypes [18]. In several samples analysed, more than one virus was found, in some cases including both DNA and RNA viruses. Taken together, this demonstrates the potential and real life utility of the microarray technique for broad-spectrum pathogen detection in clinical samples.

In 4 samples we found viruses not previously tested for in the routine analysis, HPV103 in a HPV16⁺ cervical smear sample, Picobirnavirus in a Rota A⁺ faeces sample, HAdV-C in a Sapovirus⁺ faeces sample and GBV-C in a HCV⁺ serum sample. Two of these were viruses within the same family as tested for before, HPV103 and GBV-C, while the other two were from different families. Picobirnavirus is a dsRNA virus from the Picobirnaviridae family, who was found together with the dsRNA virus Rota A (Reoviridae). HAdV-C is a dsDNA virus from the Adenoviridae family, and found together with the (+)ssRNA virus Sapovirus (Caliciviridae).

There was a false detection of virus in 3 samples, but they were all within the same family as previously tested for in the routine analysis, JCV in a BKV⁺ urine sample, Rota C in a Rota A⁺

faeces sample and Dengue 2 in a Dengue 1⁺ serum sample. They could all be explained by cross-reactivity between the probes targeting one species with other species in the same viral family. Strain or subtype identification was not a goal in the probe design, and conserved probes within a family were favoured [18]. However, future improvements to the current probe design can address cross-reactivity issues such as these described. Regarding the failure to detect Sapovirus in the double-positive Astrovirus⁺/Sapovirus⁺ faeces sample, we believe the reason was poor amplification due to low viral copy number rather than a specificity problem, since Sapovirus was detected in the Sapovirus⁺ faeces sample (Table 2). Supporting this is the Sapovirus-specific PCR used before and after Phi29-amplification (Table 1), which shows that the single-positive sample had more amplified Sapovirus (Ct=11) after amplification, compared to the double-positive sample (Ct=18) and thereby a sufficient amount for a successful detection by microarray analysis.

By microarray analysis, TTV and related viruses were detected in 77 % of the samples, from 7 different types of clinical samples. We believe this is the first report regarding the prevalence of TTV and related viruses in clinical samples from Denmark. These TTV viruses are reported to frequently infect humans, with as much as 100 % in certain populations, but no direct evidence links them to any specific clinical disease [28]. They have circular genomes (cir ssDNA) and are efficiently amplified by the Phi29 polymerase through RCA.

A microarray such as the MDA [18,19] enables simultaneous detection of viruses as well as bacteria, which could be useful in e.g. diagnosis of sexually transmitted diseases. Furthermore, many viruses cause symptoms that clinically are very similar, making it hard to choose the correct diagnostic analysis. In cases of clinical hypothesis failure, the microarray could be a useful tool for finding an etiologic agent in samples that are presumed negative. The microarray could also play a

dual role combining diagnostics and research, by being a suitable research tool for finding new pathogens.

In recent years WGA by MDA has been applied extensively, enabling researchers to get enough material from as little as a single cell for down-stream analysis [12,29,30]. Phi29 polymerase amplifies all accessible DNA present in a sample, including any exogenous DNA contamination. For viral or bacterial diagnostic purposes, where presence rather than abundance is important, the main concern is contaminating human gDNA that can compete with pathogen DNA. We show that our pre-treatment can reduce this impact from gDNA. However, in some cases this is not enough due to large amounts of gDNA being present and this, combined with low viral copy number, might make certain types of clinical samples less suitable for efficient amplification and microarray analysis. This is currently under investigation. Our microarray analysis of BKV⁺ urine samples indicated that as little as 5 genomic copies could be enough to generate enough material by Phi29-amplification to enable successful identification by the MDA. Such a high sensitivity might not be achievable for other types of viral genomes, such as linear or segmented, and is therefore under further investigation. Furthermore, to increase the chance of producing enough material from low copy number samples, we found that the WTA reaction should be allowed to run to completion (8 hours). The sample volume used for purification could also be increased.

Our results show that the protocol established for pre-treatment and amplification followed by microarray analysis is a powerful method that potentially could optimize and simplify the current way of doing multiple analyses in a diagnostic laboratory. The method detects both DNA and RNA viruses present in the same sample, regardless of size. Furthermore, the microarray analysis is shown to discriminate between different subtypes of viruses. Such an assay could also improve the

diagnostic identification of human pathogen(s) giving rise to clinical syndromes where the etiologic agent is unknown.

Author Contributions

1. Substantial contribution to conception and design, or acquisition of data, or analysis and interpretation of data: LE, MWR, CJ, KM, AF.
2. Drafting the article or revising it critically for important intellectual content: LE, MWR, AF, CJ, KM.
3. Final approval of the version to be published: LE, MWR, CJ, KM, **SG**, AF.

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FIGURE LEGENDS

Figure 1.

Comparing the effects of different pre-treatment methods on human gDNA-content (measured as β -actin) in a clinical sample. A HSV1⁺ skin lesion-sample was pre-treated with centrifugation and filtration only, or with centrifugation, filtration and DNase treatment. The β -actin content was measured by real-time PCR. (A) Centrifugation + filtration: Ct=22; centrifugation + filtration + DNase treatment: Ct=32; Δ Ct=10=1000-fold decrease in signal. (B) Time-course for Phi29-amplification of pre-treated clinical sample (centrifugation + filtration + DNase) using GenomiPhi (GE Healthcare). Fold increase in signal plotted versus amplification time for HSV1 and β -actin.

Figure 2.

WTA amplification of different clinical samples. (A) Analysis of RNA virus. HCV-specific real-time PCR on a HCV⁺ serum sample, before and after WTA. Duplicate samples. (B) Analysis of DNA virus. JCV-specific real-time PCR on a JCV⁺ CSF sample, before (undiluted) and after WTA (diluted 20-fold). (C) Purified WTA samples (HSV1, JCV and HCV) are run on a 0.8 % agarose gel to analyse size distribution, and ~500 ng per sample was loaded. Marker is TrackIt 1Kb Ladder (Invitrogen) ranging from 500 bp to 12 kb.

Figure 3.

Results from the microarray data analysis on WTA-amplified clinical samples. The lighter and darker-colored portions of the bars represent the unconditional and conditional log-odds scores, respectively. (A) Detection of HSV2 in a HSV2⁺ skin lesion-sample. (B) Detection of JCV and TTV in a JCV⁺ CSF sample. (C) Detection of HCV, GBV-C, TTV and TTV-like mini virus in a

HCV⁺ serum sample. (D) Detection of 4 different HPV subtypes (6, 16, 53 and 61) in a multi-HPV⁺ cervical smear sample.

Figure 4.

Microarray detection range for BKV⁺ clinical samples. Urine samples containing various amounts of BKV, ranging from 4.0×10^8 to 73 copies/ml (BKV1-BKV7), were Phi29-amplified (Repli-g Midi kit, Qiagen). Real-time PCR analysis using an in-house BKV-standard determined the number of BKV copies/ml before and after amplification. Selected samples were analysed by microarray. (A) Amount (copies/ml) of BKV before and after Phi29-amplification. (B) Microarray analysis on 4 selected amplified samples (BKV3-BKV6) containing various amounts of BKV.

Table 1. WTA on a diverse set of clinical samples.

Group ^a	Viral family	Virus	Size	Sample	ΔCt^b	Fold increase ^c
dsDNA	Herpesviridae	HSV1	152 kb, lin	Skin lesion	27-13	305 000
	Herpesviridae	HSV2	155 kb, lin	Skin lesion	29-15	185 000
	Papillomaviridae	HPV16	7.9 kb, cir	Cervix	26-18	24 000
	Papillomaviridae	HPV6,16,53,61 ^d	7.9 kb, cir	Cervix (6)	35-12	14 x 10 ⁶
				Cervix (16)	35-16	250 x 10 ⁶
	Polyomaviridae	BKV	5.1 kb. cir	Urine	37-16	133 x 10 ⁶
	Polyomaviridae	JCV	5.1 kb. cir	CSF	27-7	564 x 10 ⁶
dsRNA	Reoviridae	Rota A	17.4 kb, 11 seg	Faeces	12-4	330
(+)ssRNA	Astroviridae	Astrovirus	6.5 kb	Faeces	19-10	1900
	Caliciviridae	Sapovirus	7.5 kb	Faeces ^e	26-18	5800
				Faeces ^f	21-11	6100
	Flaviviridae	Dengue 1	10.7 kb	Serum	n.a.	n.a.
	Flaviviridae	HCV	9.6 kb	Serum	29-17	207 000
	Picornaviridae	HEV-A	7.4 kb	Faeces	19-6	94 000
(-)ssRNA	Paramyxoviridae	RSV	15.2 kb	TA	27-18	11 500

NOTE. HSV1, Herpes Simplex virus 1; HSV2, Herpes Simplex virus 2; HPV, Human

Papillomavirus; BKV, BK Polyomavirus; JCV, JC Polyomavirus; CSF, cerebrospinal fluid; RotaA, Rotavirus A; HCV, Hepatitis C virus; HEV-A, Human Enterovirus A; RSV, Respiratory Syncytial virus; TA, tracheal aspirate; n.a., not analyzed.

^a Viruses are grouped based on nucleic acid content, according to the Baltimore Classification.

^b Difference in Ct-value in real-time PCR before and after Phi29-amplification.

^c Fold increase after Phi29-amplification, calculated from ΔCt combined with dilution factors for each sample.

^d Only HPV6- and HPV16-specific real-time PCR's were performed.

^e Sapovirus from a double-positive Astrovirus/Sapovirus faeces sample.

^f Sapovirus from a single-positive Sapovirus faeces sample.

Table 2. Microarray analysis on a diverse set of clinical samples

Clinical sample	Type	Detected viruses	Log-Odds Score	Probe hybridization
HSV1+	Skin lesion	HSV1	1090.4	172/222
		TTV-like mini	335.2	59/107
HSV2+	Skin lesion	HSV2	977.0	171/228
HPV16+	Cervical smear	HPV16	816.6	148/374
		HPV103	541.8	58/90
		TTV	674.4	103/113
		<i>Propionibacterium phage</i>	289.1	44/50
HPV16+53+6+61+	Cervical smear	HPV6	1974.5	231/331
		HPV16	2058.2	248/375
		HPV53	2051.1	216/294
		HPV61	2082.2	231/295
BKV+	Urine	BKV	598.4	105/121
		TTV-like mini	359.7	65/107
		<i>JCV</i>	91.0	5/9
JCV+	CSF	JCV	635.1	103/125
		TTV	730.2	110/116
		<i>Phage Gifsy-2</i>	153.4	2/4
Rota A+	Faeces	Rota A	3351.9	498/696
		Picobirnavirus	677.5	49/80
		<i>Rota C</i>	284.6	35/39
		<i>Pepino mosaic virus</i>	296.2	59/126
Astrovirus+ Sapovirus+	Faeces	Astrovirus	707.9	105/120
		TTV	700.3	104/113
		<i>Lactococcus phage</i>	436.2	63/67
Sapovirus+	Faeces	Sapovirus	301.0	49/141
		HAdV-C	675.0	173/682
		TTV	858.0	199/214
		TTV midi	698.7	167/177
		TTV-like mini	699.4	175/235
		<i>Enterobacteria phage ST104</i>	224.1	63/145
Dengue 1+	Serum	Dengue type 1	719.6	92/109
		Dengue type 2	403.3	59/110
		<i>Human endogenous retrovirus</i>	333.2	26/28
HCV+	Serum	HCV	1618.0	263/268
		GBV-C	437.6	60/76
		TTV	868.7	111/112
		TTV-like mini	744.7	86/98
		<i>Phage Gifsy-2</i>	166.0	2/4
HCV+	Serum	HCV	1644.4	260/268
		GBV-C	396.8	49/76
		TTV	940.2	111/112
		TTV-like mini	799.8	85/98

		<i>Mycobacteriophage</i>	208.9	26/52
		<i>Pseudomonas phage</i>	200.3	31/66
HEV-A+	Faeces	HEV-A	1896.1	378/732
		TTV midi	638.3	145/179
		TTV	320.5	88/178
RSV+	TA	RSV	979.1	69/148
		TTV	849.0	106/112
		TTV midi	651.9	66/80
		TTV-like mini	615.3	63/84
		<i>Human endogenous retrovirus</i>	175.3	9/19
Neg control	water	n.d.	-	-

NOTE. HSV1, Herpes Simplex virus 1; TTV, Torque Teno Virus; HSV2, Herpes Simplex virus 2; HPV, Human Papillomavirus; BKV, BK Polyomavirus; JCV, JC Polyomavirus; CSF, cerebrospinal fluid; RotaA, Rotavirus A; HAdV-C, Human Adenovirus C; HCV, Hepatitis C virus; GBV-C, GB virus type C; HEV-A, Human Enterovirus A; RSV, Respiratory Syncytial virus; TA, tracheal aspirate; n.d., none detected.

Table 3. Microarray detection range for BKV⁺ urine samples

Sample	Before Phi29 copies/ml	Input per Phi29-reaction no of copies in 5µl	After Phi29 copies/ml	Detected by MDA
BKV1	4 x 10 ⁸	2 x 10 ⁶	1 x 10 ¹¹	-
BKV2	819 400	4100	5.7 x 10 ¹⁰	-
BKV3	419 700	2100	1 x 10 ¹¹	Yes
BKV4	15 810	80	1.7 x 10 ¹⁰	Yes
BKV5	892	5	1.9 x 10 ⁸	Yes
BKV6	119	0.6	11 910	No
BKV7	73	0.4	0	-